# Explicit memory creation during sleep demonstrates a causal role of place cells in navigation

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Hippocampal place cells assemblies are believed to support the cognitive map, and their reactivations during sleep are thought to be involved in spatial memory consolidation. By triggering intracranial rewarding stimulations by place cell spikes during sleep, we induced an explicit memory trace, leading to a goal-directed behavior toward the place field. This demonstrates that place cells' activity during sleep still conveys relevant spatial information and that this activity is functionally significant for navigation.

The hippocampus constitutes the neuronal substrate of the cognitive map<sup>1</sup>, as demonstrated by the firing of its principal cells in specific locations of the environment<sup>1</sup>. During sleep, hippocampal neurons replay previous wake experiences during so-called sharp wave ripples (SPW-Rs)<sup>2,3</sup>, an oscillatory pattern supporting memory consolidation<sup>4–6</sup>. However, while the animal is awake, it is impossible to dissociate its actual position from the associated neuronal firing, and the proposed function of place cells in navigation is based entirely on this correlation<sup>7,8</sup>. We hypothesized that the transient decorrelation of place cell activity from current location during sleep could provide a way to directly test the function of place cells in navigation.

In this study, we used the spontaneous activity of a given place cell during sleep to trigger automatically rewarding stimulations of the medial forebrain bundle (MFB) to create an artificial place–reward association. This version of a closed-loop-based learning procedure<sup>9–12</sup>

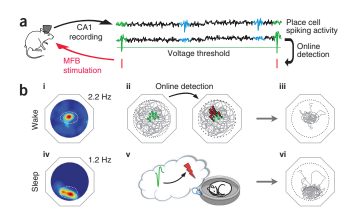
Figure 1 Pairing place cell to rewarding stimulation during wakefulness or sleep to create a place preference. (a) Method of online spike detection based on voltage threshold. Schematic waveforms from two units (blue and green) superimposed on signals recorded from hippocampal CA1 pyramidal layer (black). Spikes with high amplitude, crossing the threshold, trigger MFB stimulation. (b) Protocol for the creation of a place preference by pairing spikes to rewarding stimulations during wakefulness or sleep. (i,iv) PFs of selected pyramidal cells. (ii) During the pairing protocol performed during wakefulness, spikes of the detected place cell (green circles indicate mouse position when place cell fired) automatically trigger rewarding stimulations (red lightning bolts). (v) During sleep pairing, reactivation-related spiking activity of the selected place cell triggers rewarding stimulations. (iii,vi) Place preference is assessed by probe tests.

(**Fig. 1** and **Supplementary Fig. 1**) is similar to the classical place preference task, thought to rely on a form of hippocampus-dependent explicit memory<sup>13</sup>.

Mice were implanted with polytrodes in the CA1 pyramidal layer and stimulation electrodes in the MFB (Online Methods), known to induce reward in mice<sup>14,15</sup>. Optimal rewarding intensity was determined in a classical nose-poke procedure (Online Methods). At the intensities used, stimulations neither awakened the animals (**Supplementary Fig. 2**) nor altered basic hippocampal electrophysiological properties, such as place cell characteristics, theta oscillations or SPW-Rs (**Supplementary Figs. 3** and **4**).

As mice freely explored open field environments, spiking activity was analyzed online to determine whether a single and clearly identifiable putative pyramidal neuron displayed a place field (PF, Online Methods and **Supplementary Fig. 1**). Once selected, this place cell was used to trigger MFB rewarding stimulations (2-ms delay) during either the wake-pairing or the sleep-pairing protocol (Online Methods and **Supplementary Figs. 1** and **5**). Exploration was quantified by successive trials in PRE and POST sessions to identify exploratory bias toward the related PF.

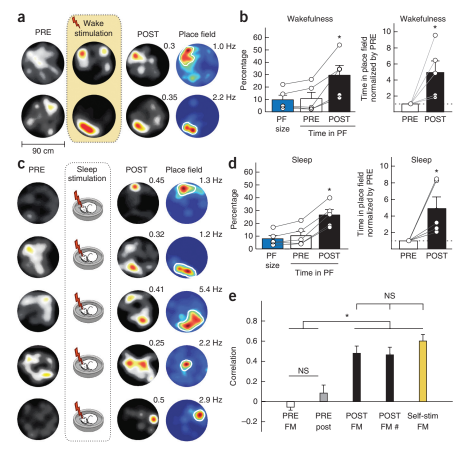
The pairing protocol was performed during wakefulness on a first set of five animals (**Fig. 2a** and **Supplementary Fig. 1a**). During the PRE session, spatial behavior was what was expected from a random and homogeneous exploration: the percentage of time mice spent in the PF corresponded to the percentage of the environment occupied by this PF (**Fig. 2b** and **Supplementary Fig. 6**). During the wakepairing session, mice stayed preferentially in the PF of the selected neuron to self-stimulate, as revealed by the high correlation between place cell firing map and occupancy map (**Fig. 2e** and **Supplementary Fig. 7**). During the POST period, time spent within the PF increased 4- to 5-fold compared to PRE, emphasizing the persistence of the memory of the place–reward association (**Fig. 2a,b**, n = 5, P = 0.043).



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Figure 2 Place preference induced after pairing place cell spikes with rewarding stimulation. (a,c) Occupancy maps before (PRE), during (a, wake stimulation) and after (POST) the session where one place cell's spikes triggered MFB stimulations, either during wakefulness (a, representative examples) or sleep ( $\mathbf{c}$ , n = 5). Color ranges from black (minimum occupancy rate) to red; peak occupancy rates are indicated above POST occupancy maps, with same scaling applied as for PRE. Corresponding firing maps (FM) of detected place cells before PRE session are represented on the right. Color ranges from blue (minimum firing rate) to red; peak firing rates indicated above. (b,d) Left, PF size expressed as a percentage of the total reachable area (blue) and percentage of time spent within the PF in PRE and POST sessions. Right, data normalized by PRE sessions after wake-pairing (b, PRE/POST, Wilcoxon matched pairs test, n = 5, Z = 2.02, \*P = 0.043) or sleep-pairing protocol (d, PRE/POST, n = 5, Z = 2.02, \*P = 0.043). (e) Correlation between occupancy maps or correlation between detected place cell's FM and occupancy map, either during PRE and POST sleep-pairing session or during wake-pairing (self-stim) sessions (Friedman test, chi-squared:  $\chi^2 = 16.60$ , n = 5, d.f. = 4, \*P = 0.0023). Correlation with FM is increased during POST (n = 5, Wilcoxon matched pairs test: correlation PRE-FM versus POST-FM, Z = 2.02, \*P = 0.043; correlation PRE-POST versus POST-FM, Z = 2.02, \*P = 0.043). Same results were observed for partial correlation (#), which quantifies



the correlation between POST and FM when the influence of PRE is removed (n = 5, correlation PRE–FM versus POST–FM given PRE, Z = 2.02, \*P = 0.043; correlation POST–FM versus POST–FM given PRE, Z = 0.73, P = 0.46). (n = 5, PRE–FM versus self-stimulation–FM, Z = 2.02, \*P = 0.043; POST–FM versus self-stimulation–FM, Z = 1.75, P = 0.07). Note the absence of correlation in PRE sessions, excluding a bias of behavior induced by arena properties. NS, not significant (PRE–FM versus PRE–POST, Z = 0.94, P = 0.34; POST–FM or POST–FM# versus self-stim–FM, Z = 1.75, P = 0.07). Error bars indicate s.e.m.

The place preference created directly using place cell activity was similar to classical place preference, induced by manual stimulations restricted to a given location (**Supplementary Fig. 8b,c**).

A second set of five animals then underwent the pairing protocol during sleep (Fig. 2c and Supplementary Fig. 1b). Analysis of the POST exploration upon awakening revealed a strong place preference for the associated PFs: the time the animals spent in these locations increased by 4- to 5-fold (Fig. 2c,d, n = 5, P = 0.043). Other behavioral measures yielded similar results (Supplementary Fig. 9).

During the 1-h sleep-pairing sessions, most of the MFB stimulations occurred during slow-wave sleep (SWS;  $567\pm80$ ) rather than rapid eye movement sleep (REM;  $11\pm7$  stimulations). Within SWS periods,  $37.4\pm8.9\%$  occurred during SPW-Rs, which represented only  $1.43\pm0.2\%$  of total sleep. This is consistent with the large increase in firing rate during SPW-Rs and with the hypothesis that SPW-Rs are essential for memory consolidation during sleep<sup>5,6</sup>. Since very few REM episodes were observed during these short sleep sessions, however, further studies would be needed to test whether this effect depends on a particular sleep stage.

Observation of animals' trajectories during POST trials of the sleeppairing protocol suggested that mice headed directly toward the PF (**Fig. 3a**). Accordingly, the time of first entry into the PF was significantly decreased in POST compared to PRE sessions. (**Fig. 3a**, n = 5, P = 0.043). All behavioral measures tended to show an increased exploration of the selected PFs during the first two trials after the sleep learning procedure when compared to the third and fourth trials, consistent with an extinction classically observed during unrewarded probe tests (**Fig. 3b**, Online Methods). Notably, the same pairing protocol during sleep with non-rewarding MFB stimulations (those that did not lead to self-stimulating behavior) did not induce any subsequent place preference upon awakening (**Fig. 3c**). Moreover, we found a significant correlation between the learning induced by the sleep-pairing protocol (an increase of time spent within the PF) and the rewarding effect of MFB stimulation (n = 7, r = 0.85, P = 0.01, **Fig. 3c**), assessed by self-stimulation in a subsequent control wakepairing session (Online Methods). Finally, there was no modification of the number of place cell–triggered MFB stimulations over time during the sleep-pairing session (**Supplementary Fig. 2e**), excluding an operant conditioning of the driving neuron, a phenomenon observed in other cellular conditioning studies<sup>9,10,12</sup>.

Taken together, these observations fulfill criteria proposed by Dickinson and Balleine to define a goal-directed strategy based on explicit knowledge of action–outcome association<sup>16</sup>. This suggests that the pairing protocol used here during sleep induced the creation of an artificial explicit memory that could be used by mice in a goal-directed way and thus goes beyond previous observations showing that certain forms of conditioning relying on a reflex response to a conditioned stimulus can take place in the sleeping or anesthetized brain<sup>17–19</sup>. Moreover, the fact that pairing spikes of a given place cell with rewarding stimulation during sleep, when its activity was decorrelated from the actual animal position, leads to a place preference toward the related place field demonstrates that place cell activity



Figure 3 Pairing MFB stimulations to place cell activity during sleep leads to a goal directed behavior upon awakening. (a) Left, firing map of the detected place cell and trial-by-trial trajectories for the POST session of the first example shown in Figure 2c. Right, quantification of the delay to reach the PF for the first time, showing a significant decrease in POST compared to PRE sessions (Wilcoxon matched pairs test, n = 5, Z = 2.02, \*P = 0.043). (**b**) Analysis comparing the first (1 and 2) and last (3 and 4) trials of the PRE and POST sessions of the sleep-pairing protocol, on time of first entry into PF (n = 5, trials 1 and 2: Z = 2.02, \*P = 0.043; trials 3 and 4: Z = 0.94, P = 0.34), cumulative distance (A.U., arbitrary units) to the PF (n = 5, trials 1 and 2: Z = 2.02, \*P = 0.043; trials 3 and 4: Z = 1.48, P = 0.14) and percentage of time spent in PF (n = 5, trials 1 and 2: Z = 2.02, \*P = 0.043; trials 3 and 4: Z = 1.75, P = 0.07). (c) Left and middle, time spent in PF in PRE (white bar) and POST (black) sessions of the sleep-pairing protocol and during self-stimulation session of the control wake protocol (green) for all mice (n = 7), including mice without self-stimulating behavior (purple dots; exclusion criterion 3). Friedman test,  $\chi^2 = 12.28$ , n = 7, d.f. = 2, P = 0.021; Wilcoxon matched pairs test, n = 7, PRE-POST sleep-pairing: Z = 2.36, \*P = 0.017; PRE-POST wake-pairing: Z = 2.36, \*P = 0.017; POST-control wake-pairing: Z = 2.19, \*P = 0.028. Right, correlation between the increase in time spent in PF during control wake self-stimulation protocol and upon awakening after sleep pairing. Spearman correlation, n = 7, r = 0.85,  $t_5 = 3.72$ , \*P = 0.01. Learning index refers to difference in percentage of time in PF, between PRE sessions and POST or during control wake self-stimulation. Error bars indicate s.e.m.

itself functions in spatial memory and navigation. Finally, our results support the theory of spatial reactivation during sleep<sup>3,4,20</sup> by showing that place cells encode the same spatial information during sleep and wakefulness.

# **METHODS**

Methods and any associated references are available in the online version of the paper.

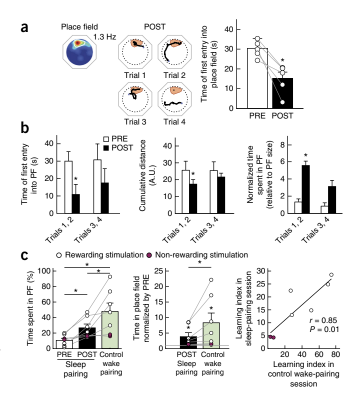
Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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# **AUTHOR CONTRIBUTIONS**

K.B. designed the experiment. G.d.L. and M.M.L. did the experiments. K.B., G.d.L. and M.M.L. analyzed the data. K.B., G.d.L., M.M.L. and L.R.-R. wrote the manuscript.



### COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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- 1. O'Keefe, J. & Nadel, L. The Hippocampus as a Cognitive Map 1-570 (Oxford University Press, 1978).
- Lee, A.K. & Wilson, M.A. Neuron 36, 1183-1194 (2002).
- Wilson, M.A. & McNaughton, B.L. Science 265, 676-679 (1994).
- Buzsáki, G. Neuroscience 31, 551-570 (1989).
- Girardeau, G., Benchenane, K., Wiener, S.I., Buzsáki, G. & Zugaro, M.B. Nat. Neurosci. 12, 1222-1223 (2009).
- Ego-Stengel, V. & Wilson, M.A. Hippocampus 20, 1-10 (2010).
- Lenck-Santini, P.-P., Muller, R.U., Save, E. & Poucet, B. J. Neurosci. 22, 9035-9047 (2002).
- Jeffery, K.J., Gilbert, A., Burton, S. & Strudwick, A. Hippocampus 13, 175-189 (2003).
- Delgado, J.M.R., Johnston, V.S., Wallace, J.D. & Bradley, R.J. Brain Res. 22, 347-362 (1970).
- 10. Shinkman, P.G., Bruce, C.J. & Pfingst, B.E. Science 184, 1194-1196 (1974).
- 11. Jackson, A., Mavoori, J. & Fetz, E.E. Nature 444, 56-60 (2006).
- 12. Fetz, E.E. Science 163, 955-958 (1969).
- 13. Eichenbaum, H. Behav. Brain Res. 103, 123-133 (1999).
- 14. Carlezon, W.A. & Chartoff, E.H. Nat. Protoc. 2, 2987-2995 (2007).
- 15. Kobayashi, T., Nishijo, H., Fukuda, M., Bures, J. & Ono, T. J. Neurophysiol. 78, 597-613 (1997).
- 16. Dickinson, A. & Balleine, B.W. Anim. Learn. Behav. 22, 1-18 (1994).
- 17. Hennevin, E. & Hars, B. Psychobiology (Austin, Tex.) 20, 166-176 (1992).
- 18. Weinberger, N.M., Gold, P.E. & Sternberg, D.B. Science 223, 605-607 (1984).
- 19. Arzi, A. et al. Nat. Neurosci. 15, 1460-1465 (2012).
- 20. Pavlides, C. & Winson, J. J. Neurosci. 9, 2907-2918 (1989).

### **ONLINE METHODS**

Subjects and surgical protocols. A total of 40 C57Bl6 male mice (Mus musculus), 3-6 months old, were implanted with movable polytrodes (six or eight twisted nichrome wires) above the CA1 pyramidal layer of right hippocampus (AP +2.2, ML +2.0, DV –1.0). Stimulation electrodes (60- $\mu$ m-diameter stainless steel) were implanted at the left medial forebrain bundle (MFB) (AP +1.4, ML +1.2, DV +4.8). During recovery from surgery (minimum 3 d) and during all experiments, mice received food and water ad libitum. The recording electrodes were progressively lowered until they reached the CA1 pyramidal layer, where ripples were recorded. Meanwhile, the mice were habituated to explore three different open fields. All mice were free of any manipulation before being included in this study. Mice were housed in an animal facility (08:00-20:00 light), one per cage after surgery. Data collection and analysis were not performed blind to the condition of the experiments, and no randomization was used. All behavioral experiments were performed in accordance with the official European guidelines for the care and use of laboratory animals (86/609/EEC) and in accordance with the Policies of the French Committee of Ethics (Decrees n° 87-848 and n° 2001–464). Animal housing facility of the laboratory where experiments were made is fully accredited by the French Direction of Veterinary Services (B-75-05-24, 18 May 2010). Animal surgeries and experimentations were authorized by the French Direction of Veterinary Services for K.B. (14-43) and L.R.-R. (75-752).

Electrophysiological procedures. *Recordings*. Signals from all electrodes were fed into a unity-gain headstage preamplifier (HS-16; Neuralynx, Bozeman, MT) and then amplified 2,000 times (tether cable to programmable amplifiers Lynx8, Neuralynx, Bozeman, MT). Signals for single-unit recordings were bandpass-filtered between 600 and 9,000 Hz. Data were digitized and stored on hard disk by a Power 1401 (CED, Cambridge, UK) acquisition system controlled by Spike2 software (CED, Cambridge, UK). Single-unit data were sampled at 20 kHz and 1.6-ms samples were time-stamped and stored for all channels of the polytrode whenever any of the channels exceeded a preset threshold. Local field potentials were sampled and stored at 1,250 Hz. During exploration or sleep recording sessions, mice were tracked by video at 30 Hz and position of the animals was reconstructed by a Matlab-based program (MathWorks, Natick, MA).

MFB stimulations. Intracranial rewarding stimulation consisted of a stimulation train lasting 100 ms and composed of 14 1-ms negative square-wave pulses (140 Hz) $^{21}$ . Optimal voltage for intracranial MFB was determined with a nose-poke task (see below). Note that these MFB stimulations did not reset theta oscillations, an effect that has been observed in some studies with stimulations located in the hypothalamic area $^{22,23}$ .

Nose-poke task. In the nose-poke task, the rodent self-administered electrical brain stimulations in an operant chamber  $(50 \times 30 \times 20 \text{ cm})$  with one panel composed of two 1.5-cm-diameter holes, in a daily session for 3 d. Each nose-poke into the left hole was detected by a photodetector and automatically triggered MFB stimulations. Mice were screened for decreasing voltage of stimulations from 8 V to 0 V in 200-s sessions. Percentage of time spent on self-stimulation was modulated by voltage intensity in accordance with previous report<sup>21</sup>. For subsequent experiments of place-reward association, we then used for each mouse the lowest stimulation intensity necessary to reach at least 80% of its time nose-poking while having no effect on sleep. Four mice did not show any self-stimulating behavior in this task and were thus excluded from the study (exclusion criterion 1).

Place preference tasks. Common paradigm. Mice were habituated to freely explore during daily 30-min sessions three distinct open fields (120 cm diameter), whose shape was either circular, square or star-shaped and surrounded by different cues. Exploration area was restricted by the length of the recording cable and all occupancy maps thus correspond only to the circular reachable area. This procedure did not affect quality of recording and was chosen to improve homogeneity of exploration as compared to that in a classical open field surrounded by walls (Supplementary Fig. 10).

Our place preference task started with a PRE session consisting of eight successive 1-min exploration trials. Then, during a pairing session, either place cell activity (sleep-pairing or wake-pairing protocols) or animal entry into a defined location (classical place preference) triggered MFB rewarding stimulations. This session was directly followed by a POST probe test, consisting of successive 1-min

exploration trials (**Supplementary Fig. 1**). As we were aiming to quantify a putative place preference, quantifications of PRE and POST sessions were done using the first four trials during which the mouse was actively exploring; trials during which mice stayed in the starting zone (within a 10-cm diameter) were not considered for the analysis. Regarding this criterion, in the sleep-pairing protocol 0.28  $\pm$  0.18 PRE trials and 1.71  $\pm$  0.28 POST trials per mouse were removed from analysis. In the wake-pairing protocol, 0.2  $\pm$  0.2 PRE trials and 0.6  $\pm$  0.4 POST trials were removed.

During PRE and POST sessions, a different starting point among four locations was assigned to each trial to force mice to use external visual cues for navigation (an allocentric strategy, known to depend on hippocampal integrity<sup>24</sup>). For each experiment, the four starting points were set symmetrically at the edge of the exploring area (**Supplementary Fig. 1d**). None of the starting points was located inside the PF, and the first trial started at the opposite side of the arena from the PF. The three remaining locations were then sequentially visited. Between trials, mice were removed from the environment and carried to the next starting point (~5-s intertrial interval).

All experiments were performed during daytime (12:00–15:00) unless specified in order to favor sleep in the sleep-pairing experiment. The behavioral task, however, was not sensitive to circadian rhythm (**Supplementary Fig. 8c–f**).

Wake-pairing protocol (using place cell activity). During the wake-pairing protocol, place cell spikes were paired with MFB stimulation at the rewarding intensity during a free exploration session, lasting 15 min after the first stimulation. The experiment was performed when spikes of a putative pyramidal cell could be efficiently and unambiguously detected online and when the corresponding PF was unique and salient (see Online Methods, "Online spike detection"). Five animals performed this wake-pairing protocol.

Sleep-pairing protocol (using place cell activity). The experiment was performed when spikes of a putative pyramidal cell reached the online spike detection criteria, and 24 of the implanted mice were therefore excluded from the study before even experiencing the sleep-pairing protocol (criterion 2). During the sleep-pairing protocol, spikes of the selected place cell were paired with MFB stimulation during 1 h of sleep. Mice slept in their home cages, which were placed outside the exploring arena and surrounded by opaque paper. An experimenter stopped the online pairing when the animal spontaneously woke up (movements seen on real-time video and decrease of the delta band in the online spectrogram). Offline analysis confirmed that the stimulation train did not modify sleep spectrograms (Supplementary Fig. 2) nor the occurrence of SPW-Rs (Supplementary Fig. 4c). Mice experienced no place preference task before the sleep-pairing protocol.

Control wake-pairing protocol (using place cell activity). The nose-pose task used to quantify the rewarding effect of MFB stimulations was done several days before the sleep-pairing protocol. It was thus important to confirm that the MFB stimulation was still rewarding at the time of the experiment. Just after the sleep-pairing protocol, efficiency of rewarding stimulations was always evaluated during a control wake-pairing protocol. These control sessions were preceded by a period of free exploration of the environment (30 min) to induce full extinction of the putative memory trace. Two mice performed the protocol with unrewarding stimulation and were therefore excluded from the main analysis (exclusion criterion 3). All other variables for these two animals were identical otherwise (Supplementary Fig. 2).

Classical place preference task (manual stimulations in an area defined by experimenter). Seven animals experienced a classical place-reward association during free exploration of the arena during light phase (12:00–15:00) and five of them experienced an ulterior session (with a different rewarded location) during dark phase (00:00–03:00). Mouse position was tracked in real time with a video camera and rewarding stimulations were delivered when mice reached a pre-established area, covering around 5% of the total surface of exploration (Supplementary Fig. 8c,d).

Exclusion criteria for behavioral experiments. As previously mentioned, 4 implanted mice were excluded because MFB stimulations did not induce self-stimulating behavior in the nose-poke task (criterion 1), 24 because they did not fulfill the criteria for online spike sorting or PF properties (criterion 2), and 2 because they did not show any self-stimulating behavior in the control wake-pairing session following the sleep-pairing protocol (criterion 3; see "Quantification of place preference at the single-mouse level").



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Offline spike sorting. Extracted waveforms were sorted via a semiautomatic cluster cutting procedure using KlustaKwik $^{25}$  and Klusters $^{26}$  (http://neurosuite.sourceforge.net/). Recordings were visualized and processed using NeuroScope and NDManager (http://neurosuite.sourceforge.net/). Isolation distance was computed with parameters proposed by Schmitzer and colleagues $^{27}$ . Since this measure depends on the number of channels, isolation measures were computed with only four of the recording channels and compared with recommended values (given for tetrode recordings). All neurons displayed isolation distance above 25 for the wake-pairing protocol and 42 for the sleep-pairing protocol, values considered as good isolation by Schmitzer and colleagues (mean isolation distance:  $86\pm23$ , n=10) $^{27}$ .

Online spike detection triggering MFB stimulations. Since the seminal work by Fetz and colleagues<sup>28–31</sup>, closed-loop systems between neural activity and either reward or electrical stimulation have been used to induce learning or to study plasticity. Yet there is still no consensus regarding the most accurate method for online spike sorting. Spikes were detected online by a simple voltage threshold applied on one specific channel of a polytrode (signal filtered above 600 Hz, referenced to a local channel without spiking activity) by a custom program in Spike2 software. This threshold was manually adjusted to detect only the highest action potentials, and the use of hexa/octrodes instead of tetrodes enhanced the probability of being in this situation. Waveforms of the detected spikes, averaged online in Spike2, allowed us to distinguish narrow spikes (usually interneuron) from large ones coming from a putative pyramidal cell. The presence of complex spikes further supported an origin from a putative pyramidal cell, and a comparison of the waveforms of all channels observed online supported detection of only one neuron (later confirmed by offline clustering). The experimenter then checked, with help of a homemade Matlab program displaying a firing rate map (FM) in real time, whether the detected putative pyramidal neuron displayed a place field. When a place cell had been detected with a given threshold, this latter was then used to automatically trigger MFB stimulation through a braincomputer interface with a delay less than 2 ms. A refractory period of 1 s from train onset was added to prevent the induced electrical artifact from being detected when crossing the threshold, leading to further stimulation not related to real spiking activity.

The quality of the online spike sorting was confirmed by comparison with offline spike sorting in terms of sensitivity and specificity (**Supplementary Fig. 5**). On average, the online spike sorting system allowed the detection of 52.9  $\pm$  9.5% of the spikes of a single pyramidal neuron as assessed by an offline spike sorting procedure, consistent with the proportion of complex spikes that do not reach the voltage threshold  $^{32}$ . During a sleep session, all spikes detected online were followed by the stimulation. Yet in spite of the electrical artifact due to stimulations, spikes were still detected by offline spike sorting, showing that  $43.2\pm10.9\%$  of spikes were followed by stimulations during the sleep-pairing protocol. Overall, this shows that the detection method is highly specific, with very few false-positive detections—our main objective—but with lower sensitivity, which was not a crucial factor for our experiment.

**Electrophysiological and behavioral analysis.** Analyses were performed with custom-made Matlab programs, based on generic code that can be downloaded at http://www.battaglia.nl/computing/ and http://fmatoolbox.sourceforge.net/.

Occupancy maps. For each exploration session, spatial density of raw trajectories in the circular reachable area was computed, normalized and smoothed. Occupancy rates thus correspond to the ratio of session time spent at a particular bin (sum of all values distributed over bins equals 1). For PRE and POST sessions composed of four trials, the same results were obtained when averaging the four maps of the successive trials, as well as when computing the map of the concatenated trajectories from those trials.

Place field analysis. A place field (PF) was defined as an area with more than nine adjacent pixels displaying a firing rate above the mean firing rate of the neuron<sup>33</sup>. The limits of the PF were then estimated by using the smoothed rate map corresponding to the area with a firing rate above 50% of the peak firing rate within the PF. To control for border effect due to our definition of the PF, behavioral analyses were also done in an increased area, referred to as "large place field," done with the Matlab function "imdilate" (Supplementary Fig. 9a), as classically done in Morris water maze experiments<sup>34</sup>.

Stability of the PF and efficiency of wake stimulations was determined by correlation analyses obtained by the bin-by-bin correlation coefficient between two different FMs or between FM and occupancy map. Partial correlation analyses were done with the function "partialcorr" in Matlab (Fig. 2e and Supplementary Fig. 7).

Behavior. Several measures were used to quantify exploration: time to enter the PF for the first time, cumulative distance defined as the instantaneous distance between the mouse position and PF center of mass, percentage of time spent within PF, and smoothed occupancy map obtained from raw trajectories. To quantify the learning effect of the pairing procedures, different normalizations were used to account for the different size of PFs: normalization by values in the PRE session or normalization by the PF size (Supplementary Fig. 9b). All comparisons between PRE and POST gave similar results when looking at ratios (POST/PRE) or differences (POST – PRE). Periods showing grooming behaviors were excluded from the analyses.

Quantification of place preference at the single-mouse level. For this quantification, we first computed the theoretical time spent in the PFs excepted by chance; that is, 10% of total exploration time spent in a PF representing 10% of the total reachable area (24 s out of 240 s corresponding to 4 trials lasting 60 s each). This was then compared to the observed proportion of time spent inside and outside the PF in the PRE session with a chi-squared test. This showed that exploration in PRE session was similar to that expected by chance. A similar chi-squared test comparison was done between PRE and POST session and between PRE and self-stimulating session in the control wake experiment. This latter analysis was used to exclude mice that did not show self-stimulating behavior in the control wake session (criterion 3).

We also computed a theoretical distribution of time spent in the PFs by using all trajectories of the PRE sessions (before any learning) taken from experiments in all protocols. As we analyzed only the first four trials in this study, one combination of four trials among the available PRE trials was assessed for each experiment. We computed the percentage of time spent in the five different PFs of the sleep protocol, for all obtained combinations except when the starting point was located within the selected PF (n = 5.914 values). This percentage was then normalized by the respective size of each PF and used to create the distribution corresponding to random exploration (**Supplementary Fig. 6c**).

Sleep scoring. To determine sleep periods, a 30-Hz camera placed 30 cm above the cage videotaped the mouse. Frame-by-frame image difference allowed determining sensitively the movements of the animal; the immobility threshold was set above respiratory movements. Sleep stages were determined by automatic k-means clustering of the theta/delta ratio extracted from the power spectrograms of hippocampal LFP signal during the episodes where the animal was immobile. REM sleep corresponded to high theta/delta ratio periods, SWS to low ratio periods.

Sharp-wave ripple detection. Offline ripple detection was performed by bandpass filtering (100–250 Hz), squaring and normalizing the field potential recorded in CA1 pyramidal layer. Ripples were defined as events peaking above 5 s.d. The beginning and end of a SPW-R were defined with a threshold of 3 s.d.

Peri-event histograms (PETHs) were calculated for the rate of spikes (in hertz) with reference to the deepest trough of all detected SPW-Rs. To compare the results between PETH centered to spike-triggered stimulation and PETH centered to SPW-Rs, a 5-ms jitter was added in the latter condition to account for the synchronization effect induced by the alignment of all ripples to the same phase, corresponding to the deepest trough (Supplementary Fig. 4g–i).

To detect the percentage of stimulation occurring during ripples, the same analysis was done on the 70 ms before the stimulation (after removing the stimulation artifact). The signal was filtered between 100 and 250Hz, squared and normalized, and we used the threshold identified by 3 s.d. obtained on stimulation-free periods. Presence of ripples was confirmed when the root mean square remained above the threshold for at least the last 30 ms before the stimulations. Note that this method is likely to underestimate the number of stimulation occurring during ripples if the spikes triggering the stimulation occur at the very beginning of the ripples.

**Statistical analysis.** Statistical analyses were done by using Matlab (Statistical Toolbox) and Statistica. Because the data did not have a normal distribution, we used non-parametric tests.

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Statistical analyses were done with Wilcoxon matched pairs tests on raw and normalized data (by PRE session values or relatively to the PF size). For comparison between different independent groups, we used the Mann-Whitney test. When multiple comparisons were present, *post hoc* group comparisons were always preceded by global analysis: Kruskal-Wallis test for unpaired protocols and Friedman test for paired protocols. In all graphics, error bars correspond to mean  $\pm$  s.e.m. and asterisks to *P* value (\*P < 0.05, \*\*P < 0.01).

Statistical power is defined as the probability of rejecting the null hypothesis when it is in fact false and should be rejected. We computed the power analysis with a Wilcoxon match paired test, comparing five values taken from a control distribution to five values taken from two different cases of test distributions. In the first situation, the control situation corresponds to five values randomly chosen according to the probability density function displayed in Supplementary Figure 6 (corresponding to performance related to five real trajectories obtained by the bootstrap procedure described in "Quantification of place preference at the single-mouse level"). Each time, those chosen values were compared to the real performance of the five mice after the sleep-pairing protocol. This was done 1,000 times and only 52 comparisons were not significant, leading to a statistical power of 0.948. In the second situation, the control situation was identical, but the test situation corresponded to a random sampling among a distribution with same mean and variance as the real performance values of the five mice after the sleep-pairing procedure. In that case, the statistical power was 0.798. Note that the same range of statistical power was obtained if we used a control distribution with the mean and variance of the real performance of our five mice during the PRE session (that is, before the sleep-pairing protocols). In this case, the statistical power obtained for the two situations were, respectively, 1 and 0.84. All the statistical powers of our analyses are in the range or above the 0.8 threshold classically used to define powerful statistics. Altogether, this shows that the statistical power of our analyses is strong enough to support the conclusion of our study.

For the evaluation of behavioral extinction of learning, we showed that all behavioral measures used (time spent within PF, cumulative distance, time to first entry) showed a statistical difference between PRE and POST session when considering the first two trials, but not for the last two trials. This observation suggests an extinction of the artificial association created earlier. Nevertheless, it is important to note that this is not a statistical proof. Indeed, the appropriate way to address this question is by testing interaction in a two-way ANOVA, and

it has been shown that other analyses can lead to erroneous interpretations<sup>35</sup>. However, our data did not satisfy criteria for parametric tests and there is no consensus concerning a non-parametric test for interaction in a two-way experimental design. Although the effect was observed for all behavioral measures tested, we decided to limit our results interpretation as being consistent with classical extinction.

**Histology.** After completion of the experiments, mice were deeply anesthetized with ketamine/xylazine solution (10% /1%). With the electrodes left *in situ*, the animals were perfused transcardially with saline ( $\sim$ 50 ml), followed by  $\sim$ 50 ml of PFA (4 g/100 mL). Brains were extracted and placed in PFA for postfixation for 24 h, transferred to PBS for at least 48 h, and then cut into 50- $\mu$ m-thick sections using a freezing microtome and mounted and stained with Cresyl violet. We then visually confirmed that recoding electrodes were implanted in the pyramidal layer of CA1 field and that stimulation electrodes reached the MFB.

A Supplementary Methods Checklist is available.

- 21. Carlezon, W.A. & Chartoff, E.H. Nat. Protoc. 2, 2987-2995 (2007).
- 22. McNaughton, N. & Sedgwick, E.M. Neuroscience 3, 629-632 (1978).
- 23. Paxinos, G. & Bindra, D. Physiol. Behav. 5, 227-231 (1970).
- 24. O'Keefe, J. & Nadel, L. *The Hippocampus as a Cognitive Map* 1–570 (Oxford University Press, 1978).
- Harris, K.D., Henze, D.A., Csicsvari, J., Hirase, H. & Buzsáki, G. *J. Neurophysiol.* 84, 401–414 (2000).
- Hazan, L., Zugaro, M.B. & Buzsáki, G. J. Neurosci. Methods 155, 207–216 (2006).
- 27. Schmitzer-Torbert, N., Jackson, J., Henze, D., Harris, K. & Redish, A.D. *Neuroscience* 131, 1–11 (2005).
- Delgado, J.M.R., Johnston, V.S., Wallace, J.D. & Bradley, R.J. Brain Res. 22, 347–362 (1970).
- 29. Shinkman, P.G., Bruce, C.J. & Pfingst, B.E. Science 184, 1194-1196 (1974).
- 30. Fetz, E.E. & Finocchio, D.V. Science 174, 431-435 (1971).
- 31. Jackson, A., Mavoori, J. & Fetz, E.E. *Nature* **444**, 56–60 (2006).
- Harris, K.D., Hirase, H., Leinekugel, X., Henze, D.A. & Buzsáki, G. Neuron 32, 141–149 (2001).
- 33. Muller, R.U., Kubie, J.L. & Ranck, J.B.J. J. Neurosci. 7, 1935-1950 (1987).
- 34. Maei, H.R., Zaslavsky, K., Teixeira, C.M. & Frankland, P.W. Front. Integr. Neurosci. 3, 4 (2009).
- Nieuwenhuis, S., Forstmann, B.U. & Wagenmakers, E. Nat. Neurosci. 14, 1105–1107 (2011).



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